

(1) Publication number: 0 684 522 A1

12

## **EUROPEAN PATENT APPLICATION**

(21) Application number: 95108090.2

(51) Int. Cl.6: G03F 7/029

(22) Date of filing: 26.05.95

(30) Priority: 27.05.94 DE 4418645

(3) Date of publication of application: 29.11.95 Bulletin 95/48

Designated Contracting States:
 AT BE CH DE DK ES FR GB IT LI NL PT SE

(1) Applicant: SUN CHEMICAL CORPORATION 222 Bridge Plaza South Fort Lee, NJ 07024 (US) 72 Inventor: Baumann, Harald Ziebikger Strasse 21 D-06846 Dessau (DE) Inventor: Timpe, Hans-Joachim Baumhofstrasse 165 D-37520 Osterode (DE) Inventor: Herting, Hans-Peter Am Schwarzenberg 21 D-37520 Osterode (DE)

(4) Representative: VOSSIUS & PARTNER Siebertstrasse 4
D-81675 München (DE)

- (54) Visible radiation sensitive composition and recording material producible therefrom.
- A visible radiation sensitive composition is described which comprises a binder, one or more polymerizable compounds containing at least one polymerizable group, and one or more dyes having an absorption range in the emission range of the radiation source, characterized in that said composition comprises as an initiator an initiator system consisting of a metallocene as a photoinitiator and an onium compound as a coinitiator. The visible radiation sensitive composition shows an increased radiation sensitivity compared to the known radiation sensitive compositions and is especially suitable for recording materials such as printing plates, which can, in particular, also be exposed by means of laser radiation in the visible range.

FP 0 684 522 A1

The invention relates to a visible radiation sensitive composition with a novel initiator system comprising a metallocene compound as a photoinitiator and an onium compound as a coinitiator. The visible radiation sensitive compositions according to the present invention are especially made use of for printing plates and in the field of photoresist technology.

Visible radiation sensitive compositions have been used for years in photopolymerizable compositions for the production of visible radiation sensitive materials, such as for example printing plates. In particular for novel techniques (i.e. exposure by means of lasers) an improved sensitivity is required, in particular within the visible range of the spectrum, in order to shorten the duration of the exposure. From an economical point of view, it is also important that low-intensity lasers can be used which are less expensive and more reliable than high-intensity lasers. Therefore, attempts have been made for quite some time to increase the sensitivity of visible radiation sensitive compositions which are to be used in photopolymerizable compositions.

It is known that the radical polymerization of ethylenically unsaturated compounds can be initiated by radiation with visible light in the presence of photoreducible dyes and reduction agents, such as for example amines (US-A-3,097,096). EP-A-122 223 discloses photoinitiators and photopolymerizable compositions which contain metallocenes. By means of such metallocenes it was possible to increase the sensitivity of the photopolymerizable layer and to thus reduce the necessary duration of exposure and the required power of the radiation source. Efforts were also made to obtain improved results by using further modified metallocenes, for example in EP-A-401 165, US-A-4,590,287, EP-A-255 486, EP-A-256 981 and US-A-5,106,722.

DE-A-40 08 815 describes a photopolymerizable composition comprising a polymerical binder, a radically polymerizable compound with at least one polymerizable group and at least one photooxidizable group in the molecule as well as a metallocene as a photoinitiator.

In order to obtain a further increase in sensitivity, an attempt was made to use the metallocene together with a coinitiator. EP-B-269 573 for example discloses liquid mixtures of photoinitiators using solutions of titanocene compounds in liquid photoinitiators of the  $\alpha$ -hydroxy and  $\alpha$ -aminoacetophenone derivative type. DE-A-38 32 032 describes a photopolymerizable mixture comprising a polymeric binder, a radically polymerizable compound with at least one polymerizable group, a photoreducible dye and a metallocene as an initiator as well as a coinitiator. The coinitiator is a trihalogenemethyl compound cleavable by radiation, which is to be used for increasing the visible radiation sensitivity. Those compounds having a triazine ring in the basic structure which carries two trihalogenemethyl groups are preferred.

DE-A-40 13 358 describes a special process for the production of printing plates or photoresists wherein metallocenes are used as photoinitiators, which is aimed at the improvement of photosensitivity.

US-A-3,717,558 relates to metallocenes of elements belonging to the subgroups of the periodic table in combination with a further photoinitiator having an activated halogene-containing group for use in photopolymerizable recording material. These initiator combinations, however, are very oxygen- and hydrolysis-sensitive and are therefore hardly suitable for the production of printing plates and photoresist materials.

It is also known to use a combination of special organometal compounds and onium salts in a hardening agent for polymerizable compositions (US-A-5,086,086). In this connection, organometal compounds are used as metallocenes, whose essential feature is that they comprise at least one metal-metal-sigma-bond, i.e. that at least two transition metal atoms are present in a complex. The curing agents of US-A-5,086,086 are not used in combination with dyes for light-induced polymerisation.

US-A-4,971,892 discloses photopolymerizable compositions which are especially suitable for printing plates and which are supposed to show a high degree of sensitivity towards visible radiation. As an initiator system for the radical polymerisation, these photopolymerizable compositions comprise an initiator selected from diaryliodonium salts, halogenated triazines and triarylsulfonium salts as well as a specific merocyanine dye.

Even though progress has been made in increasing the visible radiation sensitivity of photopolymerizable compositions, there is still a demand for compositions with further improved properties, in particular with an increased photosensitivity.

It is therefore an object of the invention to provide novel photopolymerizable compositions which show improved properties compared to the compositions known in the prior art, in particular an improved photosensitivity, and which are suitable for the production of printing plates with a high impression performance and of photoresists with a high degree of resistance against processing fluids in a cured state, which consist of a simple and thus economical composition and which have favorable thermal storage stability.

This object is achieved by a visible radiation sensitive composition which comprises as main components

- a) a binder,
- b) one or more polymerizable compounds containing at least one polymerizable group,
- c) one or more dyes having an absorption in the range of about 250 nm to about 700 nm, characterized in that it furthermore comprises

5

15

20

25

30

35

40

45

50

- d) at least one metallocene, the central atom of which is a metal atom taken from the fourth subgroup of the periodic table of the elements, and
- e) at least one onium compound.

5

20

30

40

The invention also relates to a visible radiation sensitive recording material, wherein the visible radiation sensitive composition according to the present invention is applied to a carrier.

The binders to be used are not restricted in any way and can be selected by the person skilled in the art in the known manner for the individual applications. Water-insoluble oligomers and polymers which are soluble in aqueous-alkaline media as disclosed in DE-A-40 08 815 are preferred. Especially preferred are binders comprising carboxyl groups, e.g. copolymers of (meth)acrylic acid and/or unsaturated homologues thereof such as crotonic acid, copolymers of maleic acid anhydride or the semi-esters thereof, products of the reaction of hydroxy group-containing polymers and dicarboxylic acid anhydrides, and mixtures thereof.

Polymerizable compounds, which can be used for the visible radiation sensitive compositions according to the invention are also known in the prior art and can be selected by the person skilled in the art for the desired application of the visible radiation sensitive compositions according to the present invention based on his general knowledge. Polymerizable compounds suitable for the purpose of the present invention are described e.g. in EP-A-445 624, US-A-2,760,863 and US-A-3,060,023. In particular the document EP-A-445 624 offers a helpful listing of polymerizable compounds which can be used in the visible radiation sensitive compositions of the present invention (cf. e.g. page 4, line 17 to page 7, line 14).

Most preferred are polymerizable compounds containing radically polymerizable olefinic double bonds.

The dyes which can be used in the compositions according to the invention are dyes known in this technical field which are also mentioned in EP-A-445 624. Sultable dyes are therefore in particular xanthene, benzoxanthene, benzothioxanthene, thiazine, pyronine, porphyrin or acridine dyes. Suitable xanthene and thiazine dyes are described for example in EP-A-287 817, suitable benzoxanthene and benzothioxanthene dyes are described in DE-A-20 25 291 and EP-A-321 828. A suitable porphyrin dye is e.g. hematoporphyrin and a suitable acridine dye is e.g. acriflavine chloride-hydrochloride. Examples of xanthene dyes are eosin B, eosin J, alcohol-soluble eosin, cyanosin, bengal rose, erythrosin, 2,3,7-trihydroxy-9-phenylxanthene-6-one and rhodamine-6 G. Examples of thiazine dyes are thionine, azure A and azure C. Examples of pyronine dyes are pyronine B and pyronin GY. Especially preferred are dyes of the type of triarylmethane, diarylmethane, xanthene, thioxanthene, thiazine, pyrazine, pyronine, aza[18]annulene, acridine or polymethine dyes.

The amount of the photoreducible dye is preferably in the range of 0.001 to 30 weight%, most preferably in the range of 0.01 to 20 weight%, based on the non-volatile parts of the composition.

In order to improve the sensitivity, the visible radiation sensitive composition according to the invention comprises an initiator system consisting of a metallocene which functions as a photoinitiator and an onium compound which functions as a coinitiator. Metallocenes especially suitable for the present invention are disclosed e.g. in EP-A-122 223 as being titanocenes. However, in addition to titanium as the central atom, the use of zirconium as the central atom of the metallocene is particularly preferred in the composition of the present invention.

Metallocenes which can be used in the composition according to the present invention are either commercially available, such as e.g. bis(cyclopentadienyl)-bis-[2,6-difluoro-3-(pyrr-1-yl)-phenyl]-titanium (CGI 784 of the company Ciba-Geigy), which represents an especially preferred compound for the composition of the present invention, or can be prepared according to the methods described in the prior art, e.g. in EP-A-122 223. Further metallocenes which are known photoinitiators are for example described in US-A-3,717,558, US-A-4,590,287 and US-A-5,106,722. Metallocenes of elements of subgroup IV of the periodic table, in particular compounds containing titanium and zirconium are, in addition to EP-A-122 223, also described in numerous further documents, as e.g. EP-A-119 162, EP-A-186 626, EP-A-242 330, EP-A-255 486, EP-A-256 981 and EP-A-269 573.

Especially preferred are metallocenes having a titanium or zirconium atom as the central atom and additionally having four aromatic ligands. More preferred are metallocenes wherein two ligands represent optionally substituted cyclopentadienyl groups and two ligands represent substituted phenyl groups. Most preferred is a metallocene wherein the substituted phenyl groups contain halogene atoms. Phenyl groups are also preferred which contain at least one fluorine in the o-position and which can furthermore be substituted by halogene atoms, alkyl or alkoxy groups with 1 to 4 carbon atoms and/or an optionally etherified or esterified polyoxyalkylene group. The polyoxyalkylene group generally has 1 to 6 oxyalkylene units.

Suitable onium salts are mentioned for example in US-A-5,086,086. Out of the possible onium salts mentioned therein iodonium, sulfonium, and phosphonium salts, and N-substituted N-heterocyclic onium salts or diazonium salts are preferred. Especially preferred are a diaryliodonium salt or an N-alkoxypyridinium salt. The selection of the counterion of the onium salt is not very critical. The following is a list of examples of special onium salts: 4,4'-dicumyliodonium chloride, N-methoxy- $\alpha$ -picolinium-p-toluene sulfonate, 4-methoxybenzene-

diazonium-tetrafluoro borate, 4,4'-bis-dodecylphenyliodoniumhexafluoro phosphate, 2-cyanoethyl-triphenyl-phosphonium chloride and bis-[4-diphenylsulfoniophenyl]sulfide-bis-hexafluoro phosphate (Degacure KI85 of the company Degussa).

Preferably, the composition contains

5

10

20

30

35

40

45

50

55

- a) about 5 to 90% by weight of a binder,
- b) about 5 to 90% by weight of polymerizable compounds,
- c) about 0.01 to 20% by weight of dyes,
- d) about 0.05 to 20% by weight of metallocene compounds, and
- e) about 0.05 to 20% by weight of an onium compound.

The visible radiation sensitive compositions of the present invention can be produced by means of methods known in the prior art and can then be processed further.

The compositions according to the invention are especially suitable as recording material in the production of planographic printing plates and in the field of photoresist technology. For this purpose, e.g. aluminum, steel, zinc, and copper foils, as well as plastic sheets made for example of polyethyleneterephthalate or cellulose acetate and screen printing substrates such as Perlon gauze can be used as substrates for the recording material of the present invention. In many cases, it is advantageous to subject the surface of the substrate to a mechanical or chemical pre-treatment which allows an adjustment of the adhesive properties of the layer, an improvement of the lithographical properties of the substrate surface or a decrease in the reflexion property of the substrate in the actinic range of the copy layer (antihalation).

The production of the visible radiation sensitive compositions is carried out by means of generally known methods, e.g. as described in EP-A-445 624. In this process, the layer components are taken up in a solvent and the solution or dispersion is applied to the selected substrate by means of pouring, spraying, immersing, with the help of rolls, etc. and is then dried. The selection of the light sources is not particularly restricted, for example, use can be made of tubular lamps, xenon pulse lamps, metal halogenide doped high-pressure mercury vapor lamp and carbon arc lamps. It is also possible to carry out the exposure by means of common projection or magnifying apparatuses under the light of a metal filament lamp or contact exposure by means of common light bulbs. The visible radiation sensitive compositions of the present invention and the recording materials containing the same are especially suitable for laser exposure, for example by means of argon ion, krypton ion, dye, helium, cadmium, or heliumneon lasers. It is part of the knowledge of the person skilled in the art how to carry out laser exposure.

Preferably, the visible radiation sensitive composition and thus the photopolymerizable layer is protected by a translucent and oxygen-impermeable layer which is soluble in the developer. Suitable layers consist for example of polyvinyl alcohol, polyvinylpyrrolidone, gelatine and the like. The thickness of these protection layers generally ranges from 0.1 to 10  $\mu$ m, preferably from 0.2 to 5  $\mu$ m.

It is, however, also possible to use an unsupported cover film which is removed prior to the developing of the copy layer. For this purpose for example polyester films are suitable.

The photosensitivity of the layer can be increased by briefly heating the layer to 60 to 180°C prior to the aqueous alkaline developing step. The heating is carried out after exposure and increases the crosslinking of the layer.

The developing step can be carried out by means of organic solvents, however, preferably by means of a slightly alkaline aqueous solution. During the developing step the unexposed parts of the layer are removed and the exposed regions of the copy layer remain on the carrier. Aqueous developer solutions can comprise a small part of solvents which are miscible with water. It can also comprise further known additives such as wetting agents, dyes or salts.

The examples further illustrate the invention. All the given parts represent parts by weight. All experiments were carried out under red light.

### Example 1

Electrochemically roughenend and anodically oxidized aluminum was used as the base material for the printing plates. The thickness of the oxide layer is 3.2 g/m<sup>2</sup>. The oxide layer was treated with an aqueous solution of polyvinylphosphonic acid. The thus produced base material was coated with a solution of the following composition by means of a knife coater:

4.83 parts of a terpolymer, obtained by polymerisation of 476 parts styrene, 476 parts methyl methacrylic acid

1.22 parts dipentaerythrol pentaacrylate

7.44 parts of an 80% methylethylketone solution of an urethane acrylate obtained by reacting Desmodur® N100 (of the company Bayer) and hydroxyethyl acrylate and pentaerythrol triacrylate

,

with a double bond content of 0.5 double bonds per 100 g after completion of the reaction of the isocyanate groups of the acrylic resin loncryl® 683 of the company Johnson Polymer (acid number 150)

0.1 parts ethyleosin

1.94 parts

5

10

15

20

25

35

50

55

0.35 parts 4,4'-dicumyliodonium chloride

0.25 parts bis(cyclopentadienyl)-bis-[2,6-difluoro-3-(pyrr-1-yl)-phenyl]titanium (CGI 784 of the compa-

ny Ciba-Geigy)

60 parts methylethylketone

52.5 parts methyl glycol 37.5 parts methanol

The plates were dried at 95°C in a circulation drying cabinet for 5 minutes. The gravimetrically determined weight of the dried layer was 1.8 g/m². Subsequently, an oxygen-impermeable layer with a dried layer weight of 1.7 g/m² was analogously applied by coating with a solution of the following composition:

50 parts polyvinyl alcohol (Airvol 203 of the company Airproducts; 12% remaining acetyl groups)

270 parts water

The drying was again carried out at a temperature of 95°C for 5 minutes. The thus produced plates were exposed in a vacuum printing frame by means of 4 tungsten light bulbs (200 W each) at a distance of 90 cm. A 13-step grey wedge starting at a density of 0.15 and having an increment of 0.15 served as an exposure matrix. Immediately after exposure, the plates were heated to 95°C for 1 minute in order to reinforce the terminated photopolymerisation. The plates were developed for 30 seconds in a developer of the following composition:

3.4 parts Rewopol® NLSS 28 (company REWO)
1.8 parts 2-phenoxyethanol
1.1 parts diethanol amine
1.0 parts Texapon® 842 (company Henkel)
0.6 parts Nekal® BX Paste (BASF)

0.2 parts 4-toluenesulfonic acid

91.9 parts water.

Then the developer solution is again rubbed on the surface for 30 seconds by means of a tampon and then the entire plate is rinsed with water. After this treatment, the exposed parts remain on the plate. After blackening, the plate shows the wedge steps as listed in Table I. After developing and washing with water, in order to test the printing properties of the plate, it is rubbed with an aqueous solution of 0.5% phosphoric acid and 6% gum arabic. The thus produced plate was used to produce 100 000 copies on an arc offset printing machine under usual printing conditions. The quality was good and the plates could have been copied further.

### Example 2

The substrate described in Example 1 was coated under the same conditions as in Example 1 with a layer of the following composition so that a layer weight of 1.75 g/cm² was obtained:

40 3.22 parts of the terpolymer used in Example 1 consisting of styrene, methyl methacrylate and methacrylic acid

0.81 parts dipentaerythrol pentaacrylate

4.96 parts
1.29 parts
of an 80% methylethylketone solution of the urethane acrylate used in Example 1
of the acrylic resin loncryl® 683 of the company Johnson Polymer (acid number 150)

45 0.097 parts ethyleosin

0.35 parts N-methoxy-α-picolinium-p-toluenesulfonate

0.25 parts bis(cyclopentadienyl)-bis-[2,6-difluoro-3-(pyrr-1-yl)-phenyl]titanium (CGI 784 of the compa-

ny Ciba-Geigy)

40 parts methylethylketone

35 parts methyl glycol

25 parts methanol

After the application of a polyvinylalcohol layer and developing the plate as described in Example 1, the wedge step number as shown in Table I was obtained. The printing results correspond to the ones of Example 1

### Example 3

The substrate described in Example 1 was coated under the same conditions as in Example 1 with a layer

of the following composition so that a layer weight of 1.9 g/cm<sup>2</sup> was obtained:

	3.22 parts	of the terpolymer used in Example 1 consisting of styrene, methyl methacrylate and methacrylic acid
	0.81 parts	dipentaerythrol pentaacrylate
5	4.96 parts	of an 80% methylethylketone solution of the urethane acrylate used in Example 1
	1.29 parts	of the acrylic resin loncryl® 683 of the company Johnson Polymer (acid number 150)
	0.097 parts	Rhodamine 6G
	0.35 parts	N-methoxy-pyridinium-p-toluenesulfonate
	0.25 parts	bis(cyclopentadienyl)-bis-[2,6-difluoro-3-(pyrr-1-yl)-phenyl]titanium (CGI 784 of the compa-
10		ny Ciba-Geigy)
	40 parts	methylethylketone
	35 parts	methyl glycol
	25 parts	methanol

After the application of a polyvinylalcohol layer and developing the plate as described in Example 1, the wedge step number as shown in Table I was obtained. The printing results correspond to the ones of Example 1

### Example 4

15

The substrate described in Example 1 was coated under the same conditions as in Example 1 with a layer of the following composition so that a layer weight of 1.7 g/cm² was obtained:

of the femouring composition to that a layer weight of the grown was obtained.		
	6.05 parts	of the terpolymer used in Example 1 consisting of styrene, methyl methacrylate and methacrylic acid
	1.52 parts	dipentaerythrol pentaacrylate
25	9.32 parts	of an 80% methylethylketone solution of the urethane acrylate used in Example 1
	2.4 parts	of the acrylic resin loncryl® 683 of the company Johnson Polymer (acid number 150)
	0.18 parts	ethyleosin
	0.66 parts	4-methoxbenzenediazonium-tetrafluoro borate
	0.47 parts <sup>1</sup>	bis(cyclopentadienyl)-bis-[2,6-difluoro-3-(pyrr-1-yl)-phenyl]titanium (CGI 784 of the compa-
30		ny Ciba-Geigy)
	75 parts	methylethylketone
	66 parts	methyl glycol

After the application of a polyvinylalcohol layer and developing the plate as described in Example 1, the wedge step number as shown in Table I was obtained. The printing results correspond to the ones of Example 1

## Example 5

47 parts

35

methanol

The substrate described in Example 1 was coated under the same conditions as in Example 1 with a layer of the following composition so that a layer weight of 1.85 g/cm<sup>2</sup> was obtained:

3.02 parts	of the terpolymer used in Example 1 consisting of styrene, methyl methacrylate and methacrylic acid
0.75 parts	dipentaerythrol pentaacrylate
4.65 parts	of an 80% methylethylketone solution of the urethane acrylate used in Example 1
1.2 parts	of the acrylic resin loncryl® 683 of the company Johnson Polymer (acid number 150)
0.09 parts	ethyleosin
0.45 parts	4,4'-bis-dodecylphenyliodonium-hexafluorophosphate
0.25 parts	bis(cyclopentadienyl)-bis-[2,6-difluoro-3-(pyrr-1-yl)-phenyl]titanium (CGI 784 of the compa-
	ny Ciba-Geigy)
38 parts	methylethylketone
33 parts	methyl glycol
24 parts	methanol
	0.75 parts 4.65 parts 1.2 parts 0.09 parts 0.45 parts 0.25 parts 38 parts

After the application of a polyvinylalcohol layer and developing the plate as described in Example 1, the wedge step number as shown in Table I was obtained. The printing results correspond to the ones of Example 1.

## Example 6

The substrate described in Example 1 was coated under the same conditions as in Example 1 with a layer of the following composition so that a layer weight of 1.8 g/cm² was obtained:

5	4.83 parts	of the terpolymer used in Example 1 consisting of styrene, methyl methacrylate and methacrylic acid
	1.22 parts	dipentaerythrol pentaacrylate
	7.44 parts	of an 80% methylethylketone solution of the urethane acrylate used in Example 1
	1.94 parts	of the acrylic resin loncryl® 683 of the company Johnson Polymer (acid number 150)
10	0.1 parts	Rhodamine 6G
	0.45 parts	2-cyanoethyl-triphenylphosphonium chloride
	0.25 parts	bis(cyclopentadienyl)-bis-[2,6-difluoro-3-(pyrr-1-yl)-phenyl]titanium (CGI 784 of the compa-
	•	ny Ciba-Geigy)
	60 parts	methylethylketone
15	52.5 parts	methyl glycol
	37.5 parts	methanol
		the state of department in Example 1 the

After the application of a polyvinylalcohol layer and developing the plate as described in Example 1, the wedge step number as shown in Table I was obtained. The printing results correspond to the ones of Example 1.

Example 7

20

The substrate described in Example 1 was coated under the same conditions as in Example 1 with a layer of the following composition so that a layer weight of 1.9 g/cm² was obtained:

	0	The second of th
25	4.83 parts	terpolymer 1 consisting of 10 parts styrene, 60 parts n-hexyl methacrylate and 30 parts me
		thacrylic acid
	1.22 parts	dipentaerythrol pentaacrylate
	7.44 parts	of an 80% methylethylketone solution of the urethane acrylate used in Example 1
	1.94 parts	of the acrylic resin loncryl® 683 of the company Johnson Polymer (acid number 150)
30	0.1 parts	Rhodamine B
	0.45 parts	4,4'-bis-dodecylphenyliodonium-hexafluorophosphate
	0.25 parts	dicyclopentadienyl-bis-pentafluorophenyl-zirconium
	60 parts	methylethylketone
	52.5 parts	methyl glycol
35	37.5 parts	methanol

After the application of a polyvinylalcohol layer and developing the plate as described in Example 1, the wedge step number as shown in Table I was obtained. The printing results correspond to the ones of Example 1.

### 40 Example 8

The substrate described in Example 1 was coated under the same conditions as in Example 1 with a layer of the following composition so that a layer weight of 1.9 g/cm<sup>2</sup> was obtained:

of the following composition so that a layer weight of 1.5 graff was seemed.			
	3.05 parts	methyl methacrylate copolymer ELVACITE 2670 (of the company DuPont, acid number = 74)	
45	0.81 parts	dimethylolpropane tetraacrylate	
	4.96 parts	hexa-functional urethane acrylate Actilane® 110 (of the company Akcros Chemicals)	
	1.29 parts	of the acrylic resin loncryl® 683 of the company Johnson Polymer (acid number 150)	
	0.12 parts	methyleosin	
	0.35 parts	bis[4-diphenylsulfonio)phenyl]sulfide-bis-hexafluoro phosphate (Degacure KI85, of the com-	
50	•	pany Degussa)	
	0.22 parts	bis(cyclopentadienyl)-bis-(pentafluorophenyl)titanium	
	60 parts	methylethylketone	
	52.5 parts	methyl glycol	
	37.5 parts	methanol	

After the application of a polyvinylalcohol layer and developing the plate as described in Example 1, the wedge step number as shown in Table I was obtained. The printing results correspond to the ones of Example 1.

### Example 9 (Comparative Example)

The substrate described in Example 1 was coated under the same conditions as in Example 1 with a layer of the following composition so that a layer weight of 1.8 g/cm² was obtained:

5	4.83 parts	of the terpolymer used in Example 1, consisting of styrene, methyl methacrylate and methacrylic acid
	1.22 parts	dipentaerythrol pentaacrylate
	7.44 parts	of an 80% methylethylketone solution of the urethane acrylate used in Example 1
	1.94 parts	of the acrylic resin loncryl® 683 of the company Johnson Polymer (acid number 150)
10	0.1 parts	ethyleosin
	0.45 parts	2-(4-methoxynaphth-1-yl)-4,6-bis-(trichloromethyl)-s-triazine
	0.25 parts	bis(cyclopentadienyl)-bis-[2,6-difluoro-3-pyrr-1-yl)-phenyl]titanium (CGI 784 of the company Ciba-Geigy)
	60 parts	methylethylketone
15	52.5 parts	methyl glycol
	37.5 narts	methanol

This composition corresponds to a composition of the prior art as disclosed in DE-A-38 32 032.

After the application of a polyvinylalcohol layer and developing the plate as described in Example 1, the wedge step number as shown in Table I was obtained. The number of printing operations was 90 000.

### Example 10 (Comparative Example)

20

40

The substrate described in Example 1 was coated under the same conditions as in Example 1 with a layer of the following composition so that a layer weight of 1.8 g/cm² was obtained:

25	4.83 parts	of the terpolymer used in Example 1, consisting of styrene, methyl methacrylate and methacrylic acid
	1.22 parts	dipentaerythrol pentaacrylate
	7.44 parts	of an 80% methylethylketone solution of the urethane acrylate used in Example 1
	1.94 parts	of the acrylic resin loncry® 683 of the company Johnson Polymer (acid number 150)
30	0.1 parts	ethyleosin
	0.45 parts	2,4-bis-trichloromethyl-6-(4-styrylphenyl)-s-triazine
	0.25 parts	bis(cyclopentadienyl)-bis-[2,6-difluoro-3-(pyrr-1-yl)phenyl]titanium (CGI 784 of the company
		Ciba-Geigy)
	60 parts	methylethylketone
35	52.5 parts	methyl glycol
	37.5 parts	methanol

This composition corresponds to a composition of the prior art as disclosed in DE-A-38 32 032.

After the application of a polyvinylalcohol layer and developing the plate as described in Example 1, the wedge step number as shown in Table I was obtained. The number of printing operations was 90 000.

### Example 11 (Comparative Example)

The substrate described in Example 1 was coated under the same conditions as in Example 1 with a layer of the following composition so that a layer weight of 1.8 g/cm² was obtained:

45	3.22 parts	of the terpolymer used in Example 1, consisting of styrene, methyl methacrylate and methacrylic acid
	0.81 parts	dipentaerythrol pentaacrylate
	4.96 parts	of an 80% methylethylketone solution of the urethane acrylate used in Example 1
	1.3 parts	of the acrylic resin loncryl® 683 of the company Johnson Polymer (acid number 150)
50	0.97 parts	ethyleosin
	0.20 parts	bis(cyclopentadienyl)-bis-[2,6-difluoro-3-(pyrr-1-yl)phenyl]titanium (CGI 784 of the company Ciba-Geigy)
	40 parts	methylethylketone
	35 parts	methyl glycol
55	25 parts	methanol

This composition corresponds to a composition of the prior art as disclosed in US-A-5,106,722.

After the application of a polyvinylalcohol layer and developing the plate as described in Example 1, the wedge step number as shown in Table I was obtained. The printing results correspond to the ones of Example

1.

5

## Example 12 (Comparative Example)

The substrate described in Example 1 was coated under the same conditions as in Example 1 with a layer of the following composition so that a layer weight of 1.75 g/cm² was obtained:

3.22 parts of the terpolymer used in Example 1 consisting of styrene, methyl methacrylate and methacrylic acid

0.81 parts dipentaerythrol pentaacrylate

of an 80% methylethylketone solution of the urethane acrylate used in Example 1
1.3 parts of the acrylic resin loncryl® 683 of the company Johnson Polymer (acid number 150)

0.33 parts 4,4'-dicumyliodonium chloride

0.97 parts ethyleosin

40 parts methylethylketone

15 35 parts methyl glycol 25 parts methanol

This composition essentially corresponds to a composition of the prior art as disclosed in US-A-4,971,892. After the application of a polyvinylalcohol layer and developing the plate as described in Example 1, the wedge step number as shown in Table I was obtained. The printing results correspond to the ones of Example

20 1.

25

30

35

40

## **Example 13 (Comparative Example)**

The substrate described in Example 1 was coated under the same conditions as in Example 1 with a layer of the following composition so that a layer weight of 1.85 g/cm² was obtained:

3.22 parts of the terpolymer used in Example 1, consisting of styrene, methyl methacrylate and methacrylic acid

0.81 parts dipentaerythrol pentaacrylate

4.96 parts of an 80% methylethylketone solution of the urethane acrylate used in Example 1

1.29 parts of the acrylic resin loncryl® 683 of the company Johnson Polymer (acid number 150)

0.35 parts 4,4'-dicumyliodonium chloride

0.25 parts bis(cyclopentadienyl)-bis-[2,6-difluoro-3-(pyrr-1-yl)phenyl]titanium (CGI 784 of the company

Ciba-Geigy)

40 parts methylethylketone

35 parts methyl glycol.

25 parts methanol

This composition essentially corresponds to a composition of the prior art as disclosed for example in US-A-5,086,086 or US-A-5,147,900, however, instead of the organometal compounds used in these documents, with which a production of printing plates was unsuccessful, a metallocene according to the present invention

After the application of a polyvinylalcohol layer and developing the plate as described in Example 1, the wedge step number as shown in Table I was obtained. The printing results correspond to the ones of Example

45

50

Table I

P. J. N.	5	
Example No.	Exposure (seconds)	Number of wedge steps obtained
1	30	6
2	40	5
3	30	4
4	40	5
5	30	6
6	50	5
7	40	3
8	40	4
9 (Comp. Example)	40	3
10 (Comp. Example)	120	2
11 (Comp. Example)	120	4
12 (Comp. Example)	240	4
13 (Comp. Example)	120	4

25

30

35

50

5

10

15

20

### Claims

1. Visible radiation sensitive composition, comprising

a) a binder,

- b) one or more polymerizable compounds containing at least one polymerizable group, and
- c) one or more dyes having an absorption in the range of about 250 nm to about 700 nm, characterized in that said composition comprises
  - d) at least one metallocene, the central atom of which is a metal atom taken from the fourth subgroup of the periodic table of the elements, and
  - e) at least one onium compound

as an initiator system.

- 2. Composition according to claim 1, wherein the metallocene compound comprises a titanium or zirconium atom as a central atom as well as four aromatic ligands.
  - 3. Composition according to claim 2, wherein two ligands optionally represent substituted cyclopentadienyl groups and two ligands represent substituted phenyl groups.
- Composition according to claim 3, wherein the phenyl groups of the metallocene are substituted by at least one halogen atom.
  - 5. Composition according to any one of claims 1 to 4, wherein the polymerizable compounds contain radically polymerizable olefinic double bonds.
  - 6. Composition according to any one of claims 1 to 5, wherein the dyes are of the type of triarylmethane, diarylmethane, xanthene, thioxanthene, thiazine, pyronine, pyrazine, aza[18]annulene, acridine or polymethine dyes.
- Composition according to any one of claims 1 to 6, wherein the onium compounds are iodonium, sulfonium, phosphonium, N-substituted N-heterocyclic onium salts or diazonium salts.

- 8. Composition according to claim 7, wherein the onium compound is a diaryliodonium salt or an N-alkoxy-pyridinium salt.
- Composition according to any one of claims 1 to 8, wherein the binder is an oligomer or polymer which is water-insoluble and soluble in an aqueous alkaline medium.
- 10. Composition according to any one of claims 1 to 9, comprising
  - a) 5 to 90% by weight of a binder,
  - b) 5 to 90% by weight of a polymerizable compound,
  - c) 0.01 to 20% by weight of a dye,
  - d) 0.05 to 20% by weight of a metallocene compound, and
  - e) 0.05 to 20% by weight of an onium compound.
- 11. Photosensitive recording material, wherein the composition according to any one of claims 1 to 10 is applied to a carrier.
  - 12. Photosensitive recording material according to claim 11, comprising a translucent, oxygen-impermeable layer which is soluble in the developer medium on the photopolymerizable layer.
- 20 13. Photosensitive recording material according to claim 11 or 12 whose photosensitivity was increased prior to the developing in an aqueous alkaline medium by means of a short-time heating process to 60 to 180°C.
  - 14. Use of a photosensitive composition according to any one of claims 1 to 10 for the coating of recording material.

25

5

10

30

35

40

45

50



# **EUROPEAN SEARCH REPORT**

Application Number EP 95 10 8090

		DERED TO BE RELEVAN		O ASSIDICATION OF THE	
Category	Citation of document with it of relevant pa	ndication, where appropriate, ssages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.CL6)	
A,D	EP-A-0 364 735 (HOE AKTIENGESELLSCHAFT) * claims *	CHST 25 April 1990	1-14	G03F7/029	
4	EP-A-0 207 893 (CIB	A-GEIGY AG) 7 January	1-14		
	* page 28; example	4 *			
				TECHNICAL FIELDS	
				GO3F (Int.Cl.6)	
	The present search report has been drawn up for all claims		1		
	Place of search	Date of campleties of the search	<del>' T</del>	Examiner	
	THE HAGUE	1 September 1995	່ ງ	-M. DUPART	
X : par Y : par doc	CATEGORY OF CITED DOCUMES ticularly relevant if taken alone ticularly relevant if combined with and ument of the same category shological background	E : earlier patent of  after the filing  D : document cited  L : document cited	ocument, but pub fate in the application for other reasons	dished on, or a	
O: por	p-written disdosure ermediate document		& : member of the same patent family, corresponding		

EPO FORM 1503 03.62 (POACOL)